PCT

(30) Priority Data:

08/897,744

WORLD INTELLECTUAL PROPERTY ORGANIZATION INTELLECTUAL PROPERTY ORGANIZATION INTERPRETATION OF THE PROPERTY ORGANIZATION O



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number:	WO 99/03584
	A1	(43) International Publication Date:	28 January 1999 (28.01.99)
· •	1		

US

(21) International Application Number: PCT/US98/14950

(22) International Filing Date: 20 July 1998 (20.07.98

(71) Applicant: YSI INCORPORATED [US/US]; 1700/1725 Brannum Lane, P.O. Box 279, Yellow Springs, OH 45387 (US).

21 July 1997 (21.07.97)

(72) Inventor: MOLES, Donald, R.; 2721 Cedarville-Yellow Springs Road, Cedarville, OH 45314 (US).

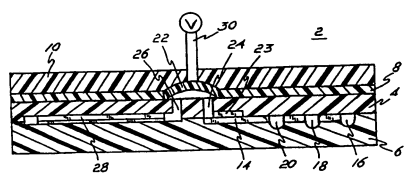
(74) Agents: LEVY, Mark, P. et al.; Thompson, Hine & Flory LLP, 2000 Courthouse Plaza, N.E., P.O. Box 8801, Dayton, OH 45401-8801 (US).

(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: MICROFLUIDIC ANALYZER MODULE



(57) Abstract

A microfluidic analyzer module. The module is constructed of a plurality of channel forming laminate layers that are directly bonded together without adhesive or other possible contaminant source located proximate the flow channels. The channel forming laminate layers define a network of fluid flow channels. A valve containing layer is directly, adhesivelessly bonded over the channel containing layers. A flexible valve member integral with the valve layer opens and closes communication between feed and sensor channels of the channel network.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL AM AT AU AZ BA BB BE BF BG BJ BR CCF CG CH CI CM CU C7, DE	Albania Armenia Austria Austria Australia Azerbaijan Bosnia and Herzegovina Barbados Belgium Burkina Faso Bulgaria Benin Brazil Belarus Canada Central African Republic Congo Switzerland Côte d'Ivoire Cameroon China Cuba Czech Republic Germany Denmark Estonia	ES FI FR GB GE GN GR HU IE IL IS IT JP KE KG KP LC LI LR	Spain Finland France Gabon United Kingdom Georgia Ghana Guinea Greece Hungary Ireland Israel Iceland Italy Japan Kenya Kyrgyzstan Democratic People's Republic of Korea Republic of Korea Republic of Korea Kazakstan Saint Lucia Liechtenstein Sri Lanka Liberia	LS LT LU LV MC MD MG MK ML MN MR MW NE NL NO NZ PL PT RO RU SD SE SG	Lesotho Lithuania Luxembourg Larvia Monaco Republic of Moldova Madagascar The former Yugoslav Republic of Macedonia Mali Mongolia Mauritania Malawi Mexico Niger Netherlands Norway New Zealand Poland Portugal Romania Russian Federation Sudan Sweden Singapore	SI SK SN SZ TD TG TJ TM TR TT UA UG US VN YU ZW	Slovenia Slovakia Senegal Swaziland Chad Togo Tajikistan Turkmenistan Turkey Trinidad and Tobago Ukraine Uganda United States of America Uzbekistan Viet Nam Yugoslavia Zimbabwe
---	--	--	---	--	---	--	--

MICROFLUIDIC ANALYZER MODULE

Field of the Invention

The present application is directed to a fluid flow module that allows for analyte sample flow therethrough while providing a situs for the location of a sensing element or elements therein to detect analyte presence and/or concentration. The module is designed for microfluidic flow rates and volumes and can be discarded after use with simple replacement by another module.

Background of the Invention

Microfluidic analyzers have been used to sense a plurality of analytes in whole blood, diluted blood, plasma or serum. Additionally in the rapidly growing fields of cell culture growth and fermentation, it is often necessary to measure glucose, glutamine, lactate, NH³, phosphate and iron in micro-scale fluid flow analysis systems.

Due to the scarce and often expensive nature of the fluid medium containing the analyte, it is imperative that the fluid medium should be conserved with only the smallest amounts used for the analysis. Also, the emphasis toward in-situ measurement of biological fluids such as cell culture media dictates that the analytical equipment should not only be small in size, but component parts of the analytical system should also be designed for easy removal from the system and rapid replacement of a new component part to the system so that cleanliness of the component can be accurately controlled without interruption in the monitoring effort. Moreover, such a component or assembly itself should be as free from contamination itself as possible.

There are several additional reasons why the measurement of sterile biological fluids benefit from the use of a disposable microfluidic; for the measurement of sterile fluids the measurement system should be enclosed, i.e. all wetted parts should be sealed so that they may be sterilized and remain that way during monitoring. If the entire fluidic system (including sensor, reagents and waste) is to be enclosed then economics dictate that it should be small, preferably very small.

Another aspect of the micro approach responds to the above-noted need to consume only small amounts of samples. If small amounts of sample are withdrawn for analysis then

small conduits are needed to convey the sample to the analyzer in order to deliver a timely result. This can be accomplished in two ways: first the conduit should be small in diameter, secondly it should be as short as possible. The best way to keep the sample conduit short is to move the analyzer to the sample. The only practical way to accomplish this is to develop the analyzer and conduit on a micro scale.

Summary of the Invention

The present invention is therefore directed to an easily assembled and manufactured microfluidic analyzer module that is adapted for easy interchangeability with a previously utilized module so that, if desired, the technician or scientist can disconnect the module and easily replace it with another. Additionally, the module itself is constructed of a plurality of directly bonded polymer layers in a sandwich laminate structure wherein the interfacial surface areas of the laminate have been etched or otherwise formed into a network of communicating microflow channels. Typically, the channels are trough-like recesses adapted for microfluid flow therein and may have widths of about .001 - .015" with channel depths being about .0005 - .015".

The module is preferably constructed of thin polymeric, laminate layers that are bonded together to form the laminate without the use of adhesives or glues that are normally used in adhesive bonding of laminate layers. This fact is especially important in critical analytical operations. Quite obviously, the presence of undesired contaminant molecules, especially those proximate the fluid containing channels, interferes with accuracy of the analytical determination.

The use of adhesives, because of the reduced scale involved, also compromises geometry. Adhesives, if liquid, tend to flow into channels, if dry, they tend not to be patternable which means that they can be mis-registered, which leads to voids along the channel or sensor cavities. These voids are "dead" volumes which destroy the desirable properties of the flow channel and make the behavior of the system less reproducible from system to system.

Further, the individual layers of the laminate can be subjected to high resolution microlithographic etching or other high resolution etching methods. When these layers are contiguously placed and bonded to each other in a sandwich construction they define a

network of small or micro dimensioned channels with certain channel portions formed in a surface area of one laminate layer and other channel portions formed in the contiguous surface area of an adjacent laminate layer. A thin, polymeric valve layer is also provided as part of the laminate structure. The valve layer is securely bonded over the sandwich construction over all surface areas except those in which valving action must occur. That is, in those areas of the sandwich in which valving action is needed to selectively block or open communication between channels of the network, an overlying flexible polymer valve is provided with flexing of the polymer valve against the underlying laminate layer surface functioning to open or block channel communication.

In accordance with the invention, complex three dimensional structures can be constructed from layers, with the benefit that each layer can be treated as a planar, two dimensional, entity. This is significant because of the extensive capabilities which have been developed which support the micromachining of planar substrates. The ability to build these layers into three dimensional structures requires not only bondability but the use of techniques of registration and bonding which preserve the integrity of the micro features themselves.

The materials used to form the module are inert, micro-machinable, bondable and dimensionally stable. Additionally, metal layers may be readily provided over these materials. This is an important consideration in regard to the desirable integration of sensor and fluidic.

The invention will be further described in conjunction with the appended drawings and following detailed description.

Brief Description of the Drawings

Fig. 1A is a plan view of a microfluidic analyzer module in accordance with the invention;

Fig. 1B is a sectional view of the microfluidic analyzer module shown in Fig. 1A taken along the plane represented by the lines and arrows 1B-1B thereof;

Fig. 1C is a sectional view of the microfluidic analyzer module similar to that shown in Fig. 1B, but here showing positioning of the valve member in the open position to allow communication between certain channels of the fluid network:

Fig. 2 is a sectional view of the microfluidic analyzer module taken along the plane represented by the lines and arrows 2-2 of Fig. 1B;

- Fig. 3 is a sectional view taken along the plane represented by the lines and arrows 3-3 of Fig. 1B;
- Fig. 3A is a magnified view of a portion of Fig. 3 showing the disposition of the sensor electrodes in the sensing channel;
- Fig. 4 is a diagrammatical section view of another microfluidic flow analyzer module taken through a plane represented by the valve layer thereof which valve layer is disposed similarly to the valve layer disposition shown in Fig. 1B;
- Fig. 5 is a schematic view of a press bonding autoclave that may be used in accordance with the invention to pressure bond layers of the microfluidic analyzer module: and
- Fig. 6 is a plan view of a platen that may be used to pressure bond the valve layer over the channel containing layers of the microfluidic analyzer module.

Detailed Description of the Preferred Embodiment

Turning now to Figs. 1A and B of the drawings, there is shown a microfluidic analyzer module 2 of the present invention comprising an upper polymer layer 4 superposed over lower polymer layer 6. As shall be explained later, the layers 4 and 6 are channel bearing layers in that they will be etched or otherwise provided with channels or portions of channels therein. Upon mating of the layers 4, 6 in superposition, a channel network is formed. A polymeric valve layer 8 is disposed on top of the layer 4 so as to selectively open and shut fluid ingress and egress channels to allow fluid flow therebetween and through the module.

As shown, lower polymer layer 6 is provided with an inlet channel 14 therein which, may be formed by suitable methods such as microlithographic etching techniques including wet, plasma, laser or e-beam etching or the like. Additionally, the channels may be formed via mechanical methods such as milling, scribing or high pressure particle stream methods. The inlet channel 14 comprises respective feed channels 16, 18, 20, in this case provided for feed of calibrant, buffer and analyte solutions to the inlet channel 14 of the analyzer module.

Upper polymer layer is provided with inlet channel 24 which is partially positioned

over channel 14. This channel communicates with perpendicularly disposed inlet channel 24 which extends perpendicularly through the layer 4 and is positioned adjacent valve egress channel 26 which, in turn communicates with sensor channel 28. The lower portion of channel 28 is formed in the lower polymer layer.

Fluid flow between the inlet channel 24 and valve egress channel 26 is selectively controlled via movement of valve member 22 which is an integral part of valve layer 8. The valve 22 surrounds the area adjacent the bores communicating with inlet and egress channels 24, 26 respectively, which bores are perpendicularly disposed with regard to and terminate in the top planar surface of upper polymer layer 4. A valve support layer 10 constructed of rigid plastic such as polyethylene, polypropylene, polycarbonate or polyphenylsulfone is provided on top of the upper polymer layer 4. The valve support layer 10 includes a vaulted concavity 12 which is positioned directly on top of the valve 22.

In one possible mode of operation a valve actuation conduit 30 is provided so that pneumatic or hydraulic fluid in communication with the conduit will selectively apply negative or positive pressure to the conduit thereby opening and closing the valve 22. In other words, valve 22 in its open position will flex so that it will be arched upwardly in the concavity 12 (see Fig. 1C) thereby allowing fluid communication between the inlet channel 24 and valve egress channel 26. Conversely, when positive pressure is applied through the conduit 30, the valve 22 will return to a position wherein it is contiguous with the top surface of upper polymer layer 4 (Fig. 1B), thereby closing communication between the inlet 24 and egress channel 26.

In the preferred method of operation, the fluid pressure will open the valve. In this way instantaneous negative pressure is not applied to the channel fluid which can cause degassing and the creation of bubbles. Bubbles are of course very undesirable in a microfluidic due to the surface tension of the bubble and the channel wall and the resulting increase in the resistance to channel flow.

It is noted that the interfacial areas defining boundaries between the upper polymer layer 4 and lower polymer layer 6 are directly bonded together without any adhesive. Also, the polymer valve layer 8 is directly bonded to the upper layer 4 without an adhesive means in all of those interfacial regions between layers 4 and 8 save for the valve areas 22.

Accordingly, considerable flex and stretch remains in the valve area 22 so that the flexible

polymer in this area may be free to flex upwardly toward the vaulted concavity 12 to open the valve and thereby provide communication between inlet 24 and egress channel 26.

Turning now to Fig. 2, there is shown lower polymer layer 6 and inlet channel 14 formed therein. Inlet channel 14 communicates respectively with calibrant branch feed channel 16, buffer feed branch channel 18, and analyte branch feed channel 20. Sensor channel 28 is in communication with valve egress channel 26.

Turning now to Fig. 3, sensor channel 28 is provided in the lower surface of upper polymer layer 4 and upper surface of layer 6. As shown in the drawing, the fluid to be analyzed passes from left to right and is in fluid flow contact with reference electrode 50, working electrode 52, and counter-electrode 54, which are formed along the sensor channel. As the fluid flows from left to right and passes over the electrodes, it flows to outlet conduit 56 and into tank 58. The outlet conduit 56 can be connected to channel 28 via snap or other quick connect mechanism to facilitate easy removal and replacement of the module 2.

As may best be seen in the magnified view, Fig. 3A, the electrodes are positioned on a plateau region 150 corresponding to the top surface of lower layer 6 within the sensor channel 28. This is accomplished by allowing a half-channel step in the sensor channel in the region of the electrodes.

It is to be remembered that although a specific electrode structure is shown in Fig. 3. other sensors could similarly be employed to contact the fluid flowing through the sensor channel. In this regard, other electrochemical or even optical sensors may be suitably disposed within sensor channel 28, in order to sense the presence or concentration of an analyte, all within the ambit of the present invention.

Turning now to Fig. 4, there is shown a sectional view of another microfluidic flow analyzer module in which a plurality of fluid flow cells are provided. Each cell is defined as including feed channel means, sensor channel means, and a valve for selectively blocking or allowing communication between the feed and sensor channels. The embodiment shown in Figs. 1A - 3 depicts a microfluidic flow analyzer module having one cell. However, in many commercial applications it will be desirable to provide a multi-cell module so that, for example, each cell may provide for analysis of a different analyte. Here, valve layer 8 includes valve areas 22a, b, and c. Underlying each area 22a, b or c are a pair of perpendicularly extending inlet and egress channel means similar to the members 24, 26

shown in Fig. 1. Each of these members terminates in a bore in the planar top surface of upper polymer layer 4. Additionally, in Fig. 4, feed conduit means have been shown for each cell and include calibrant conduits 32a, b, c, buffer conduits 34a, b, c, and analyte conduits 36a, b, c each with its associated pump, valve and reservoir. Additionally, three outlet conduits 56a, b, and c are shown, each one associated with one of the cells. A microfluidic flow analyzer module in accordance with the invention may contain a multiplicity of feed and sensor channels with each cell performing one or a variety of analyte sensing functions.

For each cell shown in Fig. 4, the feed channel 14 thereof is provided with feed conduits 32a, b or c, 34a b, or c, 36a, b or c. These conduits are snap or otherwise matingly engaged with respective branch feed channels 16, 18, 20 (such as those shown in Fig. 1A) so that the module 2 can be readily disconnected from the analytical system and replaced with another module 2. For each of the feed conduits 32a, b or c, 34a, b or c, 36a, b or c, pumps P are provided and may be operated for example by solenoid actuation through microprocessor control. The pumps include well known peristaltic pumps that are designed to pump a metered quantity of fluid from the reservoir R connected to the pumps P by valving mechanisms V. Additionally, each of the sensor channels 28 (see Fig. 3) are operatively connected to an outlet conduit 56a, b, or c via quick connect fittings. Again, this facilitates rapid replacement of one module 2 with another.

Although external pumps, valves and resevoirs are shown in the drawings and described hereinabove for purposes of clarity and simplification, it is thought that virtually all micro fluid handling processes can be accommodated within the confines of the planar interfacial surfaces of the layers 4, 6 or 4, 8. It is presently envisioned that pressurized fluid resevoirs will be used in conjunction with restrictive membranes to accomplish the desired reagent microflow.

As indicated above, any sensing means may be utilized and substituted for the electrodes 50, 52, 54. The only criterion is that the sensor, including electrochemical and optical sensors, should be capable of measuring the analyte as it flows past the sensing element in the sensor channel of the microfluidic flow analyzer module.

Additionally, multiple analysis can be conducted in one channel by in-line series disposition of multiple sensor elements along the channel. In those cases in which electrodes such as those 50, 52, 54 are shown, they may be deposited on the plateau region 150 of the

sensor channel by photo patterning, screen printing, ink jet, or micro-syringe type methods. Further, the electrodes could be covered by one or more enzyme containing membranes and interference rejection membranes if desired. In the case in which enzymes or other sensing elements are to be disposed adjacent the electrode, these can be immobilized in a polymeric medium. The polymeric medium may be electropolymerized over the electrode by, for example, the methods reported in U.S. patents 5,540,828 and 5,286,364 to Yacynych.

The microfluidic flow analyzer module in accordance with the invention can be utilized in conjunction with a plurality of different sensing methods including electrochemical amperometric, potentiometric, or conductimetric methods. Additionally, optical sensors can be utilized in conjunction with the invention although the primary concern is to provide an amperometric glucose sensor for fermentation and cell culture media. Additionally, other analytes such as glutamine, lactate, and ammonia can be sensed using the modules of the invention.

In the specific embodiment shown herein in Figs. 1A - 3, the auxiliary electrode may comprise platinum that is deposited via sputter coating, e beam evaporation or other methods. The working electrode will comprise platinum, deposited as stated above, and this electrode may be covered with an enzyme containing a laminated structure of the type known in the art. Finally, the reference electrode may comprise a platinum base over which a silver plated film is formed followed by a chloridizing treatment to form a silver/silver chloride reference electrode.

The feed and sensing channels formed in the upper and lower polymer layers of the analyzer module may be formed in the requisite surface portions of the upper and lower polymer layers by etching via photo developed micro-lithographic patterns. For example, a suitable photo resist will be applied to the substrate and then the surface of the substrate will be aligned with a mask followed by exposure to, for example, UV radiation. The etching of the polymer coating and developing of the photoresist can be accomplished in one step with a dilute aqueous alkaline solution when a positive photoresist is used. Typical etchants are dilute solutions of NaOH, KOH, tetra-alkali ammonium hydroxide, etc. The surface is then neutralized and rinsed for example with immersion in acidic acid followed by rinsing in DI water. The photoresist can be stripped with solvents such as acetone, butyl acetate or higher acetates. The skilled artisan will be able to fashion whether a positive photoresist or a

negative resist should be used and the particular types of photo resists that can be applied. Also, etching of the desired pattern may be accomplished via normal high resolution techniques including electron beam, laser or ion beam etching.

The polyimides are preferred for use as the layers 4, 6, and 8. These polymers are available commercially in thin film form with desirable film thicknesses being on the order of about 1 - 30 mils. So that the films may be bonded without application of adhesives and the like between layers, it is most desirable to employ bondable polyimides as the layers 4, 6, and 8. Suitable adhesiveless directly bondable polyimides are reported in U.S. patent 5.525,405 (Coverdall et al.). The disclosure of this patent is incorporated by reference herein.

As is reported in the '405 patent, multiple polyimide film layers may be superimposed on and directly bonded to each other without use of an adhesive. These directly bondable polyimides are aromatic polyimides containing from about 400-10.000 parts of an inorganic bonding enhancing agent such as Sn. The films comprise the polymerization - imidization product of an aromatic tetracarboxylic dianhydride component and an aromatic diamine component.

The aromatic polyimides are commonly prepared via reaction of an aromatic dianhydride and an aromatic diamine in an organic solvent medium. This results in formation of a soluble polyamic acid. This reaction normally proceeds at room temperature and is slightly exothermic. The polyamic acid precursors are converted to polyimides by dehydration, using heat or reagents such acetic anhydride or pyridine.

The most popular commercially available polyimide films are formed by the condensation reaction of pyromellitic dianhydride (PMDA) and oxydianiline (ODA). The reaction is illustrated in the following scheme:

$$(PMDA)$$

$$(ODA)$$

Ar = oxydianiline (i.e. 4,4'-diaminodiphenyl ether)

BNSDOCID +WO__9903584A1_1,+

As is stated in the '405 patent a host of other aromatic dianhydrides and aromatic diamines can be used. Suitable dianhydrides for use in the polyimide films include: pyromellitic dianhydride; 2,3,6,7-napthalene tetracarboxylic dianhydride; 3,3',4,4'-biphenyl tetracarboxylic dianhydride; 1.2,5,6-napthalene tetracarboxylic dianhydride; 2,2',3,3'-biphenyl tetracarboxylic dianhydride; 3,3',4,4'-benzophenone tetracarboxylic dianhydride; 2.2-bis(3,4-dicarboxyphenyl) propane dianhydride; 3,4,9,10-perylenetetracarboxylic dianhydride; 1,1-bis(2,3-dicarboxyphenyl) ethane dianhydride; 1,1-bis(3,4-dicarboxyphenyl) ethane dianhydride; bis(3,4-dicarboxyphenyl) methane dianhydride; bis(3,4-dicarboxyphenyl) methane dianhydride; oxydiphthalic dianhydride; bis(3,4-dicarboxyphenyl) sulfone dianhydride; and the like.

Suitable aromatic diamines for use in the polyimide films include: 4,4'-diaminodiphenyl propane: 4,4'-diamino-diphenyl methane: benzidine: 3,4'-dichlorobenzidine: 4,4'-diaminodiphenyl sulfide: 3,4'-diaminodiphenyl sulfone: 3,4'-diaminodiphenyl ether: 3,4'-diaminodiphenyl ether: 1,5-diaminonaphthalene: 4,4'-

diaminodiphenyl ethyl phosphine oxide: 4.4'-diaminodiphenyl N-methyl amine: 4.4'-diaminodiphenyl N-phenylamine: 1.4-diaminobenzene (p-phenylene diamine): 1.3-diaminobenzene; 1.2-diaminobenzene: and the like.

As mentioned in the '405 patent, copolyimides can also be prepared from the above precursors. Specifically mentioned are copolyimides derived from 15-85 mole% biphenyltetracarboxylic dianhydride. 15-85 mole% pyromellitic dianhydride. 30-10 mole% p-phenylenedramine and from 0-70 mole% 4,4'-diaminodiphenyl ether. The following copolymer is mentioned as preferred:

30 mole% pyromellitic dianhydride

20 mole% 3.3',4.4' biphenyltetracarboxylic dianhydride

30 mole% p-phenylene diamine

20 mole% 44'-diaminodiphenyl ether

The '405 patent indicates that the organic solvents include normally liquid N, N-dialkylcarboxylamides, generally. Preferred solvents include the lower molecular weight members of such carboxylamides, particularly N,N-dimethylformamide and N,N-dimethylacetamide. Other solvents which may be used dimethylsulfoxide, N-methyl-2-pyrrolidone, tetramethyl urea, dimethylsulfone, hexamethylphosphoramide, tetramethylene sulfone, and the like. The amount of solvent used preferably ranges from 75 to 90 weight % of the poly(amid acid), since this concentration has been found to give optimum molecular weight.

The solvated poly (amic) acid is further condensed to form the desired polyimide via heating at temperatures ranging from about 50°C initially, followed by higher temperature heating of about 350 - 400°C to result in almost complete condensation of the polyamic acid intermediate to the imide form.

So as to enhance the adhesive-less bonding properties of the films a metallic bond enhancing agent such as Sn is added to the polyimide during the polymerization or film forming process. As reported in the '405 patent, Sn in the amount of about 400 -10.000 ppm has proven effective. Reputedly, the preferred amount of Sn ranges from 1,000 - 4,000 ppm with the most preferred amount being about 1,200 - 3,500 ppm. Tin can be added as an organotin compound as described in U.S. patent 5,272,194 or a Sn⁻² or Sn⁻⁴ salt as specified

in U.S. patent 5.218,034. Bis-tributyltinoxide is mentioned as a preferred organotin compound.

In accordance with the invention, the layers 4, and 6 of the microfluidic flow analyzer module are preferably composed of DuPont® XNATM Kapton® polyimide film. This is available and advertised as an adhesiveless, self-bondable film. The preferred film thickness is 5 mils.

With regard to the valve layer 8 in accordance with the invention, preliminary investigation has indicated that this can be composed of DuPont's Kapton® KJ thermoplastic polyimide film. The preferred thickness for this film is about 1 mil.

After the layers 4 and 6 have been purchased or prepared, the lower surface of layer 4 and top surface of layer 6 are etched in accordance with conventional techniques, as stated above, to provide the required pattern which will correspond to the sensor and feed channels in the finished microfluidic flow analyzer module. The perpendicularly arranged (i.e. with respect to the surface plane) channels 24, 26 can be inserted via conventional techniques including wet, plasma, E-beam, laser drilling or similar operations. The sensor means, such as electrodes, etc. can be placed along one of the layer surfaces defining a portion of the sensor channel 28. The layers 4, 6 are then placed in superposed relation with the etched surfaces thereof in mating, interfacial engagement to define the fluid flow channels needed for fluid inlet and outlet from the feed channel through the sensor channel.

As illustrated diagrammatically in Fig. 5, the layers 4, 6 are then placed in a vacuum autoclave 100 or similar hot press apparatus. As shown, upper platen 102 is placed on top of the layers 4.6 in the autoclave with lower platen 104 on the bottom. The platens may be hydraulically driven together to form a pressure nip on the layers 4.6.

The bonding operation may be carried out in the autoclave at temperatures of around 350 - 455°C at pressures of about 24 - 690 bar for a period of about 5 minutes to three hours. Preferably, the heat-press bonding operation is carried out under vacuum of less than 760 mm Hg. In this manner the sandwich construction providing the channel bearing layers 4, 6 is formed.

After the sandwich construction of layers 4, and 6 is provided, the valve layer 8 can be laminated over the top of the sandwich construction, again in a vacuum autoclave as shown in Fig. 5 or in another hot press. Here, bonding to the sandwich is normally carried out at about

275 - 350°C at about 200 psi under vacuum conditions.

Notably, when bonding the layer 8 to the sandwich assembly 4, 6, relief areas or apertures 122 are provided at least in the top platen 102 (as is depicted in Fig. 6). This platen (as well as the platens used in forming the original sandwich) has length and width dimensions that roughly match the dimensions of the layers 4,6, 8. However, during the bonding of the valve layer 8 to the sandwich, the relief areas 122 overlie the valve areas 22 that will appear in the finished module. During the pressing operation, less pressure will be exerted on the laminate in the areas 122 than along the other laminate areas under the solid portions of the platen. Accordingly, the layer 8 in those areas will not readily bond to layer 4 in the valve areas. The layer 8 in those areas will remain moveable and flexible so that it will perform as a valve that upon actuation will selectively open and close communication between inlet channel 24 and valve egress channel 26. Additionally, the relief areas may be provided with vents so as to prevent build up of gas pressure from outgass products.

After the laminate consisting of layers 4, 6, and 8 is provided, the valve support layer 10 is bonded thereover by conventional means such as by use of a dry sheet adhesive or heat bonding of thin elastomer sheet in a liquid solvent medium. The microfluidic analyzer module comprising layers 4, 6, 8 and 10 can then, if desired, be mounted on a base or other mounting mechanism ready for insertion as a component of a fluid analysis system. The base may be attached/sealed to the module 2 using conventional means such as self-sticking double sided adhesive, heat-bondable sheet adhesive or through the use of an elastomeric gasket seal in combination with a glue or solvent applied to the perimeter of the rigid base and receiving structure.

The preceding description and accompanying drawings are intended to be illustrative of the invention and not limited. Various other modifications and applications will be apparent to one skilled in the art without departing from the true spirit and scope of the invention as defined by the following claims.

What is claimed is:

1. A module with fluid flow channels formed therein, comprising a first and second polymer layer, said first layer superposed over said second layer to define an interfacial area with micro-fluid channels formed along said interfacial area, said first and second polymer layers bonded directly together along said interfacial area without adhesive, said micro-fluid channels having widthwise dimensions of from about .001 - .015" and depths of about from .0005 - .015".

- 2. A module as recited in claim 1 wherein said fluid channels comprise a fluid inlet channel and a fluid sensing channel, and valve means operatively associated with said fluid inlet and fluid sensor channels for selectively opening and closing communication between said fluid inlet channel and said fluid sensing channel.
- 3. A module as recited in claim 2 wherein an analyte sensor is operatively associated with said fluid sensing channel.
- 4. A microfluidic analyzer module comprising:
 - (a) a first sheet of polyimide having a top surface area and a bottom surface;
 - (b) a second sheet of polyimide material having a top surface area and a bottom surface, said first and second polyimide sheets in superposed relation with said bottom surface of said first sheet of polyimide and said top surface of said second sheet of polyimide forming bonded, adhesiveless, interfacial surface areas, said interfacial surface areas comprising fluid flow channels therein.
- 5. A microfluidic module as recited in claim 4 wherein said flow channels comprise a feed channel and a sensor channel, said module further comprising valve means operatively associated with said feed channel and said sensor channel for selectively opening and closing communication between said feed channel and said sensing channel, and a sensing element disposed in said sensor channel to detect the presence or concentration of an analyte in said sensor channel.

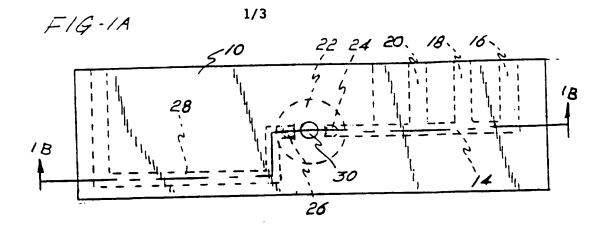
6. A microfluidic analyzer module as recited in claim 5 wherein said feed channel comprises a feed channel bore extending upwardly through said first sheet of polyimide and terminating in said top surface of said first sheet of polyimide. said sensor channel comprising a sensor channel bore extending upwardly through said first sheet of polyimide and terminating in said top surface of said first sheet of polyimide proximate said feed channel bore, said valve means comprising a flexible material positioned over said feed channel bore and said sensor channel bore, actuation means connected to said valve means for pressing said flexible material over said feed channel bore and said sensor channel bore to block communication therebetween and for relaxing said flexible material in spaced relation over said top surface of said first sheet of polyimide for opening communication therebetween.

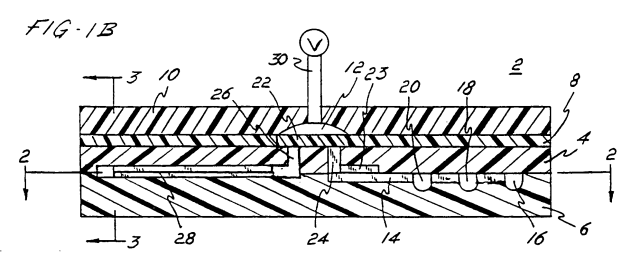
- 7. A microfluidic analyzer module as recited in claim 6 wherein said sensing element comprises an electrochemical sensing element.
- 8. A microfluidic analyzer module as recited in claim 6 wherein said sensing element comprises an optical sensing element.
- 9. A microfluidic analyzer module as recited in claim 7 wherein said electrochemical sensing element comprises an electrode.
- 10. Laminated microfluidic analyzer assembly module comprising a first, second and third layer of polymeric material, said first and second layers directly bonded together without adhesive to form a sandwich structure having a top surface and a bottom surface, said sandwich structure comprising a fluid flow channel formed therein, said fluid flow channel being in communication with said top surface of said sandwich structure, a valve region of said top surface in operative association with said fluid flow channel, said third layer of polymeric material superposed over said top surface of said sandwich structure, a portion of said third polymeric material overlying said valve region to define a valve, said third layer of polymeric material comprising a flexible plastic material bonded over said top surface of said sandwich structure except for said valve, said third layer of polymeric material retaining its flexibility at said valve, and a valve actuator operatively associated with said valve for

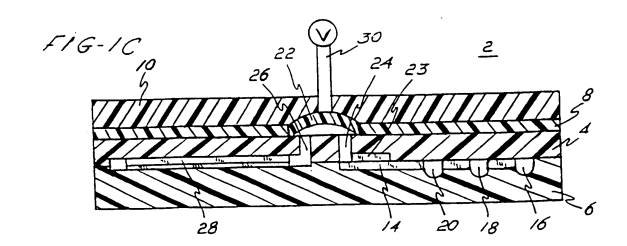
selectively closing and opening said valve.

11. Laminated microfluidic analyzer module as recited in claim 10 wherein said third layer is directly, non-adhesively bonded to said top surface of said sandwich structure.

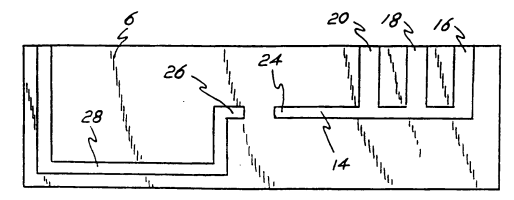
- 12. Laminated microfluidic analyzer module as recited in claim 11 wherein said first, second and third layers of polymeric material include polyimide polymers.
- 13. Laminated microfluidic analyzer module as recited in claim 10 further including a fourth layer overlying said third layer, said fourth layer comprising a concave surface portion overlying said valve in said third layer.
- 14. Laminated microfluidic analyzer module as recited in claim 13 wherein said valve actuator comprises a pressure source communicating with said concave surface portion.



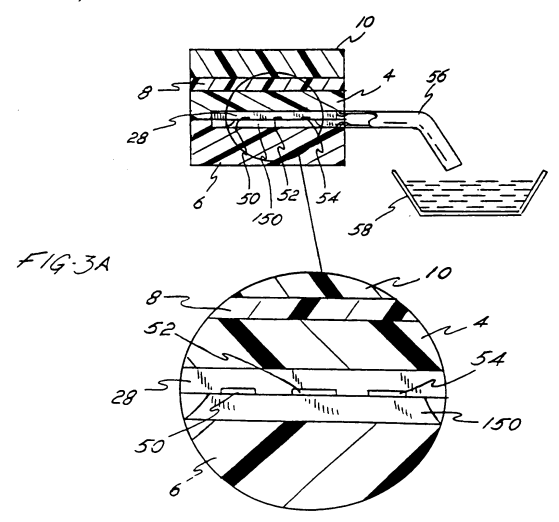


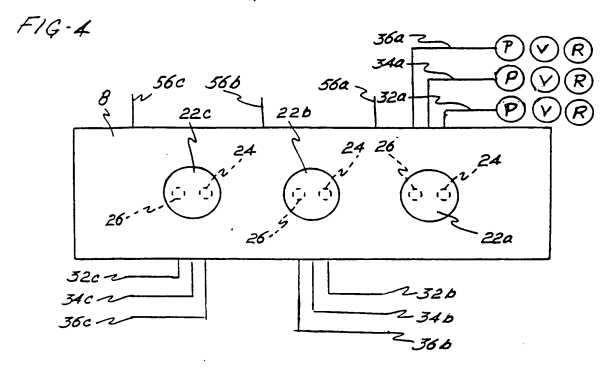


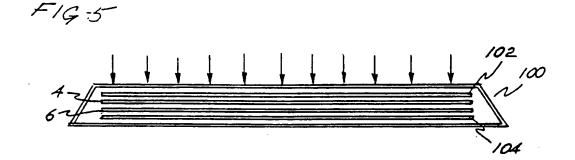


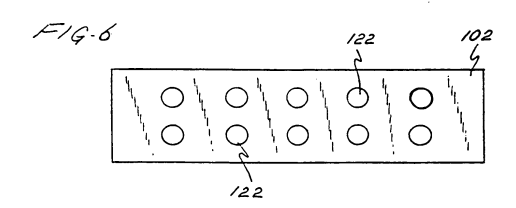


F1G-3









A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 B01L3/00 B32B27/34

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum occumentation searched (classification system followed by classification symbols) IPC 6-801L

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

x	EP 0 770 871 A (HEWLETT PACKARD CO) 2 May 1997	Relevant to claim No
	2 May 1997	1
3	· · · · · · · · · · · · · · · · · · ·	
	see column 6, line 55 - column 7, line 16 see column 11, line 30 - line 40 see column 9, line 53 - column 10, line 7	1,4
x	see column 12, line 11 - line 26; figure 1	1,4
Y	see column 12, line 44 - column 13, line 38	2,3,5-9
	see column 16, line 37 - column 17, line 8; figures 6.7	
Y	WO 95 08716 A (ROSEMOUNT ANALYTICAL INC) 30 March 1995	2,3.5-9
	see page 5, line 20 - page 8, line 20; figures	
	see page 10, line 23 - page 11. line 11	
	-/	

X Patent family members are listed in annex
"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"Y" document of particular relevance, the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of mailing of the international search report 05/11/1998
Authorized officer Hocquet . A

1

INTERNATIONAL SEARCH REPORT

PCT/US 98/14950

State of document with indication where appropriate is the reservant cassages Perevant to claim No 2.5.10		US 98/14950
26 September 1989 see column 4, line 54 - column 5, line 3 see column 6, line 25 - line 35 X W0 97 02357 A (AFFYMETRIX INC :ANDERSON ROLFE C (US); LIPSHUTZ ROBERT J (US); RAV) 23 January 1997 see page 32, line 24 - page 33, line 38 see page 34, line 31 - page 35, line 9 see page 36, line 1 - line 7 see page 36, line 34 - line 32; figure 2B x see page 8, line 27 - line 31 see column 27 - line 31 see column 9, line 55 - line 31 see column 9, line 58 - column 10, line 5 EP 0 394 048 A (BIOTRACK INC) 24 October 1990 see column 5, line 7 - line 29; figures EP 0 716 918 A (DU PONT) 19 June 1996 cited in the application see page 1, line 15 - line 25; claim 1 A W0 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3, line 22 - line 30 see page 5. line 3 see page 34, line 10 - line 29 see page 34, line 15 - line 28 W0 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24, line 20 - page 27, line 10;		Relevant to claim No
ROLFE C (US); LIPSHUTZ ROBERT J (US); RAV) 23 January 1997 see page 32, line 24 - page 33, line 38 see page 34, line 31 - page 35, line 9 see page 36, line 1 - line 7 see page 36, line 34 - line 32; figure 2B X see page 8, line 27 - line 31 Z5 see page 8, line 27 - line 31 Z6 December 1995 see column 2, line 25 - line 31 see column 9, line 58 - column 10, line 5 A EP 0 394 048 A (BIOTRACK INC) 24 October 1990 see column 5, line 7 - line 29; figures A EP 0 716 918 A (DU PONT) 19 June 1996 cited in the application see page 1, line 15 - line 25; claim 1 A WO 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29 see page 34, line 15 - line 28 A WO 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24, line 20 - page 27, line 10;	26 September 1989 see column 4, line 54 - column 5, line 3	2.5.10
X see page 36, line 34 - line 32; figure 2B	ROLFE C (US); LIPSHUTZ ROBERT J (US); RAV) 23 January 1997 see page 32, line 24 - page 33, line 38 see page 34, line 31 - page 35, line 9	1-9
26 December 1995 see column 2, line 25 - line 31 see column 9, line 58 - column 10, line 5 A EP 0 394 048 A (BIOTRACK INC) 24 October 1990 see column 5, line 7 - line 29; figures A EP 0 716 918 A (DU PONT) 19 June 1996 cited in the application see page 1, line 15 - line 25; claim 1 A W0 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29 see page 34, line 15 - line 28 A W0 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24. line 20 - page 27, line 10;	see page 36, line 34 - line 32; figure 2B	
24 October 1990 see column 5, line 7 - line 29; figures EP 0 716 918 A (DU PONT) 19 June 1996 cited in the application see page 1, line 15 - line 25; claim 1 W0 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29 see page 34, line 15 - line 28 A W0 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24. line 20 - page 27, line 10;	26 December 1995 see column 2, line 25 - line 31	1
cited in the application see page 1, line 15 - line 25; claim 1 WO 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29 see page 34, line 15 - line 28 WO 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24. line 20 - page 27, line 10;	24 October 1990	1,4,10
29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29 see page 34, line 15 - line 28 A WO 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24. line 20 - page 27, line 10;	cited in the application	1,4,10
12 June 1997 see page 24, line 20 - page 27, line 10;	29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30. line 10 - line 29	1,4,10
	12 June 1997 see page 24, line 20 - page 27, line 10;	13
		US 4 869 282 A (SITTLER FRED C ET AL) 26 September 1989 see column 4, line 54 - column 5, line 3 see column 6, line 25 - line 35 WO 97 02357 A (AFFYMETRIX INC :ANDERSON ROLFE C (US); LIPSHUTZ ROBERT J (US); RAV) 23 January 1997 see page 32, line 24 - page 33, line 38 see page 34, line 31 - page 35, line 9 see page 36, line 1 - line 7 see page 36, line 34 - line 32; figure 2B see page 8, line 27 - line 31 US 5 478 751 A (OOSTA GARY M ET AL) 26 December 1995 see column 2, line 25 - line 31 see column 9, line 58 - column 10, line 5 EP 0 394 048 A (BIOTRACK INC) 24 October 1990 see column 5, line 7 - line 29; figures EP 0 716 918 A (DU PONT) 19 June 1996 cited in the application see page 1, line 15 - line 25; claim 1 WO 94 21386 A (RES CORP TECHNOLOGIES INC) 29 September 1994 see page 3. line 22 - line 30 see page 5. line 3 see page 30, line 10 - line 29 see page 30, line 10 - line 29 see page 30, line 15 - line 28 WO 97 21090 A (GAMERA BIOSCIENCE) 12 June 1997 see page 24, line 20 - page 27, line 10; figures 6-11

Form PCT (SA210 (continuation of second shaet) (July 1992)

1

INTERNATIONAL SEARCH REPORT

information on patent family members

PCT/US 98/14950

	nt document i search report		Publication date	1	Patent ramily member(s)	Publication date
EP 0	770871	A	02-05-1997	US	5641400 A	24-06-1997
WO 9	508716	Α	30-03-1995	CA CN	2169826 A 1133080 A	30-03-1995 09-10-1996
				JP	9505130 T	20-05-1997
US 4	869282	Α	26-09-1989	CA	2004965 A	09-06-1990
				DE	68921892 D	27-04-1995
				DE	68921892 T	23-11-1995
				EP	0447451 A	25-09-1991
				WO	9006470 A	14-06-1990
WO 9	9702357	Α	23-01-1997	AU	6404996 A	05-02-1997
				EP	0843734 A	27-05-1998
US S	 5478751	Α	26-12-1995	AU	1446195 A	17-07-1995
				CA	2178505 A	06-07-1995
				EΡ	0737105 A	16-10-1996
				JP	9507572 T	
				WO	9517966 A	06-07-1995
EP (0394048	Α	24-10-1990	US	4952373 A	
				AT	112509 T	
				AU	618461 B	
				AU	5249490 A	
				CA	1334737 A	
				CA	2013695 A	
				DE	69013043 D	
				DE	69013043 T	
				ES	2064623 T	
				JP	2561364 B	
				JP 	3041361 A	21-02-1991
ΕP	0716918	Α	19-06-1996	US	5525405 A	
				JP 	8224845 A	03-09-1996
WO	9421386	Α	29-09-1994	US	5686548 A	
				US	5686549 A	11-11-1997

INTERNATIONAL SEARCH REPORT

Application No PCT/US 98/14950

Patent document cited in search report		Publication date		atent family member(s)	^D ublication date
WO 9721090	A		EP NO AU WO	0865606 A 982563 A 4144897 A 9807019 A	23-09-1998 05-08-1998 06-03-1998 19-02-1998

Form PCT/ISA 210 (patent tamely annex) (July 1992)

